CASE REPORT

Brucellosis as a trigger agent for Henoch-Schönlein purpura

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Abstract: Vasculitis in childhood is a result of a spectrum of causes ranging from idiopathic conditions with primary vessel inflammation to syndromes after exposure to recognized antigenic triggers, such as infectious agents and drugs causing hypersensitivity reactions. Henoch–Schönlein purpura (HSP) is the most common vasculitis of childhood. Although there is often a history of a recent or simultaneous upper respiratory tract infection, no consistent causative organism is found. We report an 11-year old boy with HSP and brucellosis and we speculated that brucellosis was the trigger agent for HSP (Ref. 13). Full Text in PDF www.elis.sk.

Key words: brucellosis, trigger agent, Henoch-Schönlein purpura, vasculitis, infectious agents, drug, hypersensitivity reactions, primary vessel inflammation.

Hence-Schönlein purpura (HSP) is characterized by non-thrombocytopenic purpura, arthritis and arthralgia, abdominal pain and gastrointestinal hemorrhage, and glomerulonephritis (1). It is one of the most common vasculitides of childhood (2). HSP was first recognized by Heberden in 1801 and first described as an association between purpura and arthritis by Schönlein in 1837. In 1874 and in 1899, Henoch added descriptions of gastrointestinal involvement and renal involvement, respectively (3, 4). The diagnostic criteria of HSP were modified by the European League against Rheumatism/ Paediatric Rheumatology European Society (EULAR/PReS) as follows (5).

Classification criteria for Henoch–Schönlein purpura

Palpable purpura (mandatory criterion) in the presence of at least one of the following four features:
- Diffuse abdominal pain
- Any biopsy showing predominant IgA deposition
- Arthritis (Acute, any joint) or arthralgia
- Renal involvement (any haematuria and/or proteinuria)

Many reports have implicated infections such as parvovirus B19, hepatitis B virus, hepatitis C virus, human immunodeficiency virus, Streptococcus species particularly β-hemolytic streptococci, Salmonella species, Shigella species, Staphylococcus aureus, medications such as antibiotics, angiotensin converting enzyme inhibitors, non-steroidal anti-inflammatory (NSAI) agents and toxins such as vaccinations, insect bites and food allergy as a potential trigger for this disease. (6, 7) In this study, we report an 11-year old boy with HSP and brucellosis as a potential trigger infection.

Case report

The 11-year old boy was referred from orthopedics department to our clinic because of arthritis, fever, weight loss, abdominal pain, and purpura skin lesion of the lower legs. His past history was unremarkable except for consumption of fresh unpasteurized cheese. Physical examination revealed a blood pressure of 120/85 mmHg, bilateral symmetric palpable purpura on the lower extremities, abdominal defense-tenderness and swelling on the bilateral ankle joints. On admission, laboratory investigations revealed hemoglobin 11.8 g/dl, white blood cell count 9900/μL, platelet count 319,000/μL, C-reactive protein 19.1 mg/dl, blood urea nitrogen 7 mg/dl, and serum creatinine 0.68 mg/dl. Serum electrolyte, liver enzymes, blood coagulation tests, antinuclear antibody, rheumatoid factor, antistreptolysin O, serum IgA and urinalysis were within normal levels. The brucella agglutination test was ≥1/1280. On skin punch biopsy, capillaries and venules of the dermis were affected by a leukocytoclastic vaskulitis with vessel wall necrosis and perivascular accumulation of polymorphonuclear leukocytes and mononuclear cells. Immunofluorescence microscopy was positive for IgA.

The patient was hospitalized with the diagnosis of HSP and brucellosis; antihistaminic and NSAI drugs were initiated for pruritis and arthritis, respectively and rifampycin and tetracycline were initiated for brucellosis. On the 2nd day of admission, the patient was discharged with these medications. Two weeks after discharging from the hospital, he was readmitted to our pediatric nephrology outpatient clinic for follow up and his purpura on the lower extremities was extinguished and the swelling on the bilateral ankle joints was relieved. Antihistaminic and NSAI drugs were discontinued and rifampycin and tetracycline were continued
totally for six weeks. On the 3rd month of follow-up, he was fully recovered without any systemic involvement of HSP.

Discussion

HSP is defined as vasculitis with predominantly IgA deposits in the walls of small vessels in the skin, gastrointestinal tract and kidney associated with arthralgias or arthritis (8). The etiology is unknown, but HSP follows an upper respiratory infection (9). Infection agents, some medications and some toxins were reported as a trigger for HSP (7). Zucchini et al reported two cases of HSP associated with acute salmonella enterocolitis (10). Courtney PA et al reported a case of vasculitis occurring 7 days after meningitis C vaccination in a 17 year old girl (11). In different studies, hepatitis A and B, Mycoplasma pneumonia, Bartonella henselae and Parvovirus B19 have been implicated as a trigger for HSP (1).

Human brucellosis, caused by organisms of the genus Brucella, is still a major public health problem worldwide. Brucella abortus (cattle), B. melitensis (goat/sheep), B. suis (swine), and B. canis (dog) are the most common organisms responsible for human disease (12). It is a systemic illness that can be very difficult to diagnose in children without a history of animal or food exposure. Our patient had a history of consumption of fresh unpasteurized cheese. In the diagnosis of brucellosis, the routine laboratory investigations of the blood generally are not helpful; instead the serum agglutination test is the most widely used and detects antibodies against B. abortus, B. melitensis, and B. suis. In our patient the brucella agglutination test was $\geq 1/1280$ and we speculated that brucella was a trigger infection for HSP. Massasso et al, reported a case of HSP with brucellosis and they concluded the ability of brucellosis to mimic systemic vasculitic disease, in particular HSP (13). In their study, they showed the leukocytoclastic vasculitis with IgA immunofluorescence of dermal vessels consistent with HSP, so we thought, both in our case and in theirs, brucellosis was the trigger agent for HSP. In conclusion, we would like to show that our case suggests the possibility of brucellosis being a trigger agent of HSP.

References


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